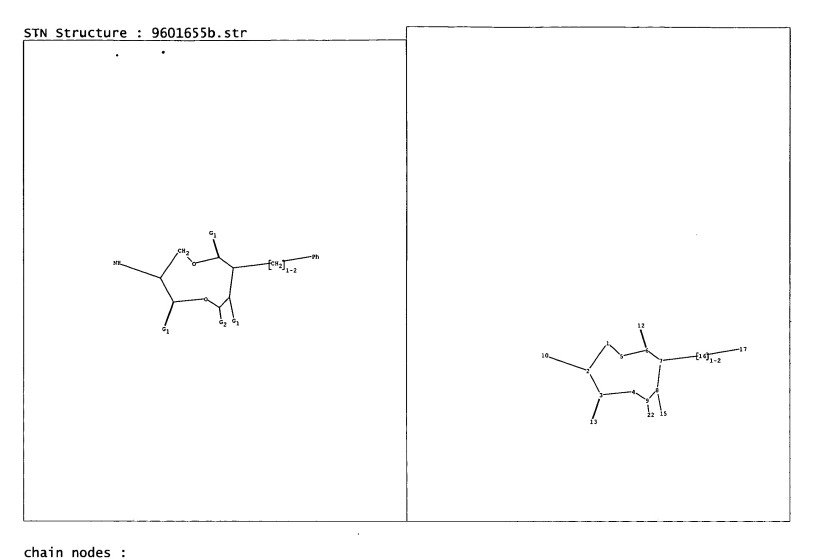
=> d his

(FILE 'HOME' ENTERED AT 11:46:22 ON 09 NOV 2001) FILE 'REGISTRY' ENTERED AT 11:46:25 ON 09 NOV 2001 L1STRUCTURE UPLOADED L2 14 S L1 L3 231 S L1 FULL FILE 'HCAPLUS' ENTERED AT 11:47:11 ON 09 NOV 2001 L417 S L3 L5 10 S L4 AND PD < MARCH 1998 1 S L5 AND SAKANAKA, O?/AU L6 9 S L5 NOT L6 L7 O S L7, IBIB ABS FHITSTR, 1-9 L8 FILE 'CAOLD' ENTERED AT 11:50:40 ON 09 NOV 2001 => s 13L9 0 L3



```
10 12 13 15 16 17 22

ring nodes:
    1 2 3 4 5 6 7 8 9

chain bonds:
    2-10 3-13 6-12 7-16 8-15 9-22 16-17

ring bonds:
    1-2 1-5 2-3 3-4 4-9 5-6 6-7 7-8 8-9

exact/norm bonds:
    2-10 3-13 6-12 8-15 9-22

exact bonds:
    1-2 1-5 2-3 3-4 4-9 5-6 6-7 7-8 7-16 8-9 16-17

isolated ring systems:
    containing 1:
```

G2:CH3,Et

Match level:

G1:0,5

Match level:
1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:Atom 8:Atom 9:Atom 10:CLASS 12:CLASS 13:CLASS 15:CLASS 16:CLASS 17:Atom 22:CLASS

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                 The CA Lexicon available in the CAPLUS and CA files
         Feb 06
NEWS
                 Engineering Information Encompass files have new names
         Feb 16
NEWS
                 TOXLINE no longer being updated
         Apr 23
NEWS
                 Search Derwent WPINDEX by chemical structure
NEWS
         Apr 23
                 PRE-1967 REFERENCES NOW SEARCHABLE IN CAPLUS AND CA
         May 07
                 DGENE Reload
NEWS
      7
         Jun 20
NEWS
      8
                 Published patent applications (A1) are now in USPATFULL
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         JUL 13
                 New SDI alert frequency now available in Derwent's
                 DWPI and DPCI
NEWS 10
         Aug 23
                 In-process records and more frequent updates now in
                 MEDLINE
NEWS 11
         Aug 23
                 PAGE IMAGES FOR 1947-1966 RECORDS IN CAPLUS AND CA
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         Aug 23
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NEWS 13
         Sep 17
                 IMSworld Pharmaceutical Company Directory name change
                 to PHARMASEARCH
NEWS 14
         Oct 09
                 Korean abstracts now included in Derwent World Patents
                 Index
         Oct 09
NEWS 15
                 Number of Derwent World Patents Index updates increased
NEWS 16
         Oct 15
                 Calculated properties now in the REGISTRY/ZREGISTRY File
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         Oct 22
                 Over 1 million reactions added to CASREACT
NEWS 18
         Oct 22
                 DGENE GETSIM has been improved
        Oct 29
NEWS 19
                 AAASD no longer available
             August 15 CURRENT WINDOWS VERSION IS V6.0c,
NEWS EXPRESS
              CURRENT MACINTOSH VERSION IS V6.0 (ENG) AND V6.0J (JP),
              AND CURRENT DISCOVER FILE IS DATED 07 AUGUST 2001
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COST IN U.S. DOLLARS

SINCE FILE TOTAL ENTRY SESSION 0.15 0.15

FULL ESTIMATED COST

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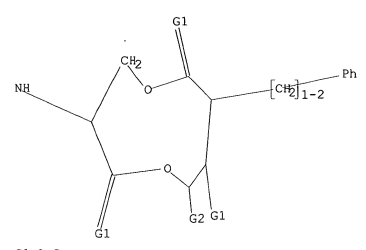
Calculated physical property data is now available. See HELP PROPERTIES for more information. See STNote 27, Searching Properties in the CAS Registry File, for complete details: http://www.cas.org/ONLINE/STN/STNOTES/stnotes27.pdf

=> Uploading 9601655b.str

L1 STRUCTURE UPLOADED

=> d l1

L1 HAS NO ANSWERS L1 STR



G1 O,S G2 Me,Et

Structure attributes must be viewed using STN Express query preparation.

=> s 11

SAMPLE SEARCH INITIATED 11:46:55 FILE 'REGISTRY'
SAMPLE SCREEN SEARCH COMPLETED - 15 TO ITERATE

100.0% PROCESSED 15 ITERATIONS 14 ANSWERS

SEARCH TIME: 00.00.02

FULL FILE PROJECTIONS: ONLINE **COMPLETE**

PROJECTED ITERATIONS: 68 TO 532

PROJECTED ANSWERS: 56 TO 504

L2 14 SEA SSS SAM L1

=> s l1 full

THE ESTIMATED SEARCH COST FOR FILE 'REGISTRY' IS 133.25 U.S. DOLLARS DO YOU WANT TO CONTINUE WITH THIS REQUEST? (Y)/N or END:y

FULL SEARCH INITIATED 11:47:06 FILE 'REGISTRY'
FULL SCREEN SEARCH COMPLETED - 241 TO ITERATE

100.0% PROCESSED 241 ITERATIONS 231 ANSWERS SEARCH TIME: 00.00.02

L3 231 SEA SSS FUL L1

=> file hcaplus

COST IN U.S. DOLLARS SINCE FILE TOTAL

Page 3

ENTRY SESSION 133.56 133.71

FULL ESTIMATED COST

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FILE COVERS 1947 - 9 Nov 2001 VOL 135 ISS 20 FILE LAST UPDATED: 7 Nov 2001 (20011107/ED)

This file contains CAS Registry Numbers for easy and accurate substance identification.

This file supports REG1stRY for direct browsing and searching of all substance data from the REGISTRY file. Enter HELP FIRST for more information.

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=> s 13

17 L3 L4

=> s 14 and pd < march 1998

17342633 PD < MARCH 1998 (PD<19980300)

L5 10 L4 AND PD < MARCH 1998

=> s 15 and sakanaka, o?/au

22 SAKANAKA, O?/AU L6 1 L5 AND SAKANAKA, O?/AU

=> d l6, ibib abs fhitstr, 1

ANSWER 1 OF 1 HCAPLUS COPYRIGHT 2001 ACS 1999:19692 HCAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER:

130:168617

TITLE:

UK-2A, B, C and D, novel antifungal antibiotics from Streptomyces sp. 517-02 III. Absolute configuration

of

an antifungal antibiotic, UK-2A, and consideration of

its conformation

AUTHOR(S): Shibata, Kozo; Hanafi, Muhammad; Fujii, Jyunko;

Sakanaka, Osamu; Iinuma, Katsuharu; Ueki,

Masashi; Taniguchi, Makoto

CORPORATE SOURCE: Faculty of Science, Osaka City University, Osaka,

558-8585, Japan

SOURCE: J. Antibiot. (1998), 51(12), 1113-1116

CODEN: JANTAJ; ISSN: 0021-8820

PUBLISHER: Japan Antibiotics Research Association

DOCUMENT TYPE: Journal LANGUAGE: English

GΙ

AB The abs. configuration of UK-2A (I) was detd. by the elucidation of the abs. configurations of butanolide II and the serine deriv. III, the products of alk. hydrolysis of I. The abs. configuration of UK-2A was found to be (+)-(2R,3R,4S,7S).

IT **167173-86-6**, UK 2B

RL: MSC (Miscellaneous)

(detn. of the abs. configuration of UK-2A, an antifungal antibiotic)

RN 167173-86-6 HCAPLUS

CN 2-Butenoic acid, 2-methyl-, (3S,6S,7R,8R)-3-[[(3-hydroxy-4-methoxy-2-

pyridinyl)carbonyl]amino]-6-methyl-4,9-dioxo-8-(phenylmethyl)-1,5-dioxonan-7-yl ester, (2E)- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (+). Double bond geometry as shown.

REFERENCE COUNT: REFERENCE(S):

5

(1) Fuji, K; Synthesis 1975, P276 HCAPLUS

(2) Hanafi, M; J Antibiotics 1996, V49, P1226 HCAPLUS

(3) Trecourt, F; Tetrahedron 1993, V49, P8373 HCAPLUS

(4) Ueki, M; J Antibitics 1996, V49, P639 HCAPLUS

(5) Wasserman, H; Chem Rev 1986, V86, P845 HCAPLUS

ALL CITATIONS AVAILABLE IN THE RE FORMAT

=> d his

(FILE 'HOME' ENTERED AT 11:46:22 ON 09 NOV 2001)

FILE 'REGISTRY' ENTERED AT 11:46:25 ON 09 NOV 2001

L1 STRUCTURE UPLOADED

L2 14 S L1

L3 231 S L1 FULL

FILE 'HCAPLUS' ENTERED AT 11:47:11 ON 09 NOV 2001

L4 17 S L3

L5 10 S L4 AND PD < MARCH 1998

L6 1 S L5 AND SAKANAKA, O?/AU

=> s 15 not 16

L7 9 L5 NOT L6

=> s 17, ibib abs fhitstr, 1-9

1332 'L7'

14 IBIB

179901 ABS

4 ABSES

179905 ABS

(ABS OR ABSES)

O FHITSTR

6636974 1

1404846 9

L8 0 L7, IBIB ABS FHITSTR, 1-9

('L7'(W)IBIB(W)ABS(W)FHITSTR(W)1(W)9)

=> d 17, ibib abs fhitstr, 1-9

L7 ANSWER 1 OF 9 HCAPLUS COPYRIGHT 2001 ACS ACCESSION NUMBER: 1999:313243 HCAPLUS

DOCUMENT NUMBER: 131:214101

TITLE: Total synthesis of the antifungal dilactone UK-2A and

analogs and their bioactivities

AUTHOR(S): Kamei, Noriyuki; Shibata, Tetsuo; Inoguchi, Kiyoshi;

Senda, Hisato; Ikari, Takashi; Itoh, Nobuko; Shimano,

Masanao

CORPORATE SOURCE: Department of Medical Chemistry and Molecular Design,

Drug Discovery Research Laboratories, Kaken

Pharmaceutical Co., Ltd., Japan

SOURCE: Tennen Yuki Kagobutsu Toronkai Koen Yoshishu (

1998), 40th, 679-684

CODEN: TYKYDS

PUBLISHER: Nippon Kagakkai

DOCUMENT TYPE: Journal LANGUAGE: Japanese

GI

AB UK-2A (I) which has recently been isolated from the mycelial cake of Streptomyces sp. 517-02, possesses nine-membered dilactone and a picolinic

II

acid moiety. The plane structure of UK-2A has been elucidated by 1H and 13C NMR analyses and chem. degrdn. studies, but the relative and abs. configurations of the four chiral centers in UK-2A still remain to be detd. UK-2A has strongly inhibited the growth of various kinds of yeasts and filamentous fungi, but its cytotoxic activities against several kinds of mammalian cells were very weak. The combination of its interesting mol. architecture and the potent antifungal activity prompted us to

Ι

launch

the total synthesis of UK-2A. The synthesis of UK-2A has been achieved

through a 12-step sequence from II in 26% overall yield. The key strategy employed in this approach involves; (1) construction of the three

consecutive chiral centers from C2 to C4 based upon the well-established Evans aldol reaction and (2) the nine-membered lactonization. The authors' synthetic route to UK-2A would permit a practical and reliable construction of UK-2A and a variety of its analogs. In order to define the selective cytotoxicities of UK-2A against yeasts and filamentous fungi, UK-2A and its analogs synthesized were subjected to the MIC evaluation and cytotoxic activity examn. compared with the ref. compds., amphotericin B and fluconazole. UK-2A has a broad antifungal spectrum, while its cytotoxicities was considerably weak compared to other substrates. The results of the UK-2A analogs suggested that the basicity of the picolinic acid moiety in UK-2A was essential for the antifungal activities and that the feature of the nine-membered dilactone contributed

to the selective cytotoxicities.

167173-85-5P, Antibiotic UK 2A

RL: BAC (Biological activity or effector, except adverse); PRP (Properties); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)

(total synthesis of antifungal dilactone UK-2A and analogs and bioactivities)

RN 167173-85-5

ANSWER 2 OF 9 HCAPLUS COPYRIGHT 2001 ACS L7 1998:651994 HCAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER: 130:3703

TITLE: Total synthesis of the antifungal dilactones UK-2A

and

UK-3A: the determination of their relative and absolute configurations, analog synthesis and

antifungal activities

Shimano, Masanao; Kamei, Noriyuki; Shibata, Tetsuo; AUTHOR(S):

Inoguchi, Kiyoshi; Itoh, Nobuko; Ikari, Takashi;

Senda, Hisato

CORPORATE SOURCE:

Dep. Med. Chem. Mol. Design, Drug Discovery Res.

Lab.,

SOURCE:

Kaken Pharmaceutical Co., Ltd., Minami Kawara-cho,

Yamashina-ku, Kyoto, 607-8042, Japan Tetrahedron (1998), 54(42), 12745-12774

CODEN: TETRAB; ISSN: 0040-4020

PUBLISHER: DOCUMENT TYPE:

Journal

LANGUAGE:

English

OTHER SOURCE(S):

CASREACT 130:3703

Elsevier Science Ltd.

GI

The synthesis of the antifungal dilactones (I), UK-2A (R = OMe) and UK-3AAΒ (R = H), is described. In addn. to providing a workable synthetic route to these potent antifungal antibiotics, this has allowed us to det. the assignment of the relative and abs. configurations in the nine-membered ring. Furthermore, UK-2A analogs were also synthesized and evaluated for their antifungal activities and cytotoxic activities along with UK-2A, (2R, 3R, 4S, 7R)-UK-2A, UK-3A, (2R, 3R, 4S, 7R)-UK-3A, and antimycin A. The structural requirements for the selective cytotoxicity against yeasts and filamentous fungi will also be suggested.

167173-85-5P, UK-2A ΙT

> RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)

(synthesis, antifungal activity, cytotoxicity and abs. configuration

of

dilactones UK-2A and UK-3A)

167173-85-5

REFERENCE COUNT:

REFERENCE(S):

(2) Barrow, C; J Antibiot 1997, V50, P729 HCAPLUS (3) Brooks, B; J Comput Chem 1983, V4, P187 HCAPLUS

(4) Centeno, N; Chem Phys Lett 1995, V232, P374 **HCAPLUS**

(5) Dickie, J; J Med Chem 1963, V6, P424 HCAPLUS

(6) Evans, D; J Am Chem Soc 1981, V103, P2127 HCAPLUS

ALL CITATIONS AVAILABLE IN THE RE FORMAT

HCAPLUS COPYRIGHT 2001 ACS ANSWER 3 OF 9

ACCESSION NUMBER:

CORPORATE SOURCE:

1998:355895 HCAPLUS

DOCUMENT NUMBER:

129:122477

TITLE:

Enantioselective total synthesis of the antifungal dilactone, UK-2A: the determination of the relative

and absolute configurations

AUTHOR(S):

Shimano, Masanao; Shibata, Tetsuo; Kamei, Noriyuki

Dep. Medicinal Chem. Molecular Design, Drug Discovery Res. Labs., Kaken Pharmaceutical Co., Kyoto,

607-8042,

SOURCE:

Tetrahedron Lett. (1998), 39(24), 4363-4366

CODEN: TELEAY; ISSN: 0040-4039

PUBLISHER:

Elsevier Science Ltd.

DOCUMENT TYPE:

Journal English

LANGUAGE: OTHER SOURCE(S):

CASREACT 129:122477

GΙ

AB The synthesis of the antifungal dilactone, UK-2A (I), is described. In addn. to providing a workable synthetic route to this potent antifungal antibiotic, this has allowed us to det. the assignment of the relative

and

abs. configurations in the nine-membered ring.

IT 167173-85-5P, (+)-UK-2A

RL: PRP (Properties); SPN (Synthetic preparation); PREP (Preparation) (abs. configuration of UK-2A via enantioselective total synthesis)

RN 167173-85-5

L7 ANSWER 4 OF 9 HCAPLUS COPYRIGHT 2001 ACS ACCESSION NUMBER: 1998:22846 HCAPLUS

DOCUMENT NUMBER:

128:163891

TITLE:

The mode of action of UK-2A and UK-3A, novel

antifungal antibiotics from Streptomyces sp. 517-02

AUTHOR(S):

Ueki, Masashi; Taniguchi, Makoto

CORPORATE SOURCE:

Dep. Biology, Fac. Sci., Osaka City Univ., Osaka,

558,

Japan

SOURCE:

J. Antibiot. (1997), 50(12), 1052-1057

CODEN: JANTAJ; ISSN: 0021-8820

PUBLISHER:

Japan Antibiotics Research Association

DOCUMENT TYPE: Journal LANGUAGE: English

AB UK-2A and UK-3A are structural relatives of antimycins, which were isolated as antifungal antibiotics with little cytotoxicity that demonstrated inhibition of respiratory activity. They halve the cellular respiration of yeast within 4~5 min and the intracellular ATP content within 2~5 min. They inhibited the yeast mitochondrial respiration using .beta.-hydroxybutyrate and succinate as a respiratory substrate, but no inhibition was obsd. using ascorbate-reduced tetra-Me p-phenylenediamine as the substrate. The site of respiratory inhibition of UK-2A and UK-3A was thought to be the cytochrome bcl complex in the mitochondrial

electron

transport chain of yeast cells. They also inhibited the mitochondrial respiration of rat liver. Intact animal cells might have some system to defend themselves from the actions of UK-2A and UK-3A.

IT **167173-85-5**, UK-2A

RL: BUU (Biological use, unclassified); BIOL (Biological study); USES (Uses)

(mechanism of antifungal action of UK-2A and UK-3A)

RN 167173-85-5

L7 ANSWER 5 OF 9 HCAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER:

1997:504110 HCAPLUS

DOCUMENT NUMBER:

127:217524

TITLE:

UK-3A, a novel antifungal antibiotic from

Streptomyces

sp. 517-02: fermentation, isolation, structural

elucidation and biological properties

AUTHOR(S):

Ueki, Masashi; Kusumoto, Atsushi; Hanafi, Muhammad; Shibata, Kozo; Tanaka, Toshio; Taniguchi, Makoto Faculty of Science, Osaka City University, Osaka,

CORPORATE SOURCE:

558,

Japan

SOURCE:

J. Antibiot. (1997), 50(7), 551-555

CODEN: JANTAJ; ISSN: 0021-8820

PUBLISHER:

Japan Antibiotics Research Association

DOCUMENT TYPE:

Journal

LANGUAGE:

English

GI

AB A novel antifungal antibiotic, UK-3A (I), was obtained from the mycelial cake of Streptomyces sp. 517-02. I was very similar in structure to UK-2A, a structural relative of antimycin A. The antifungal spectrum of I

was relatively broad (MICs for yeasts and filamentous fungi: 1.56.apprx.6.25 and 0.39.apprx.1.56 .mu.g/mL, resp.). The cytotoxic activity of I was weak (IC50: 18.apprx.100 .mu.g/mL).

IT 194931-82-3P, Antibiotic UK 3A

RL: BAC (Biological activity or effector, except adverse); BOC (Biological $\ensuremath{\mathsf{BOC}}$

occurrence); BPN (Biosynthetic preparation); PRP (Properties); BIOL (Biological study); OCCU (Occurrence); PREP (Preparation) (UK-3A is a novel antifungal antibiotic from Streptomyces)

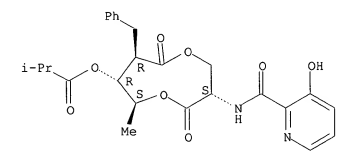
RN 194931-82-3 HCAPLUS

CN Propanoic acid, 2-methyl-, (3S,6S,7R,8R)-3-[[(3-hydroxy-2-

pyridinyl)carbonyl]amino]-6-methyl-4,9-dioxo-8-(phenylmethyl)-1,5-dioxonage 11

7-yl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (+).



ANSWER 6 OF 9 HCAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER:

1997:16443 HCAPLUS

DOCUMENT NUMBER:

126:144017

TITLE:

UK-2A, B, C and D, novel antifungal antibiotics from

Streptomyces sp. 517-02. II. Structural elucidation

AUTHOR(S):

Hanafi, Muhammad; Shibata, Kozo; Ueki, Masashi;

Taniguchi, Makoto

CORPORATE SOURCE:

Fac. Sci., Osaka City Univ., Osaka, 558, Japan

SOURCE:

J. Antibiot. (1996), 49(12), 1226-1231

CODEN: JANTAJ; ISSN: 0021-8820

PUBLISHER:

Japan Antibiotics Research Association

DOCUMENT TYPE:

Journal

English LANGUAGE: UK-2A, UK-2B, UK-2C and UK-2D, novel antibiotics produced by Streptomyces AΒ

sp. 517-02, exhibit strong antifungal activity. The structures were elucidated based on spectral and chem. evidence that these compds. are

the

derivs. of the nine-membered dilactone formed from serine and 4-hydroxypentanoic acid moiety.

IT167173-86-6P

> RL: PRP (Properties); PUR (Purification or recovery); PREP (Preparation) (structural elucidation of UK-2A, UK-2B, UK-2C and UK-2D, novel antifungal antibiotics from Streptomyces sp. 517-02)

167173-86-6 HCAPLUS RN

CN 2-Butenoic acid, 2-methyl-, (3S,6S,7R,8R)-3-[[(3-hydroxy-4-methoxy-2-

pyridinyl)carbonyl]amino]-6-methyl-4,9-dioxo-8-(phenylmethyl)-1,5-dioxonan-7-yl ester, (2E)- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (+). Double bond geometry as shown.

L7 ANSWER 7 OF 9 HCAPLUS COPYRIGHT 2001 ACS ACCESSION NUMBER: 1996:463922 HCAPLUS

DOCUMENT NUMBER: 125:109869

TITLE: UK-2A, B, C and D, novel antifungal antibiotics from

Streptomyces sp. 517-02. I. Fermentation, isolation,

and biological properties

AUTHOR(S): Ueki, Masahi; Abe, Keiichi; Hanafi, Muhammad;

Shibata,

Kozo; Tanaka, Toshio; Taniguchi, Makoto

CORPORATE SOURCE: Fac. Science, Osaka City Univ., Osaka, 558, Japan

SOURCE: J. Antibiot. (1996), 49(7), 639-643

CODEN: JANTAJ; ISSN: 0021-8820

DOCUMENT TYPE: LANGUAGE: Journal English

GΙ

Α.

AB Novel antifungal antibiotics, UK-2A (I), UK-2B (II) and a mixt. of UK-2C and UK-2D, were obtained from the mycelial cake of Streptomyces sp. 517-02. All of the UK-2 compds. were similar in structure to antimycin

The antifungal activities of of UK-2 compds. were as strong as that of antimycin A. However, the UK-2 compds. demonstrated weak cytotoxicity compared to antimycin A.

IT 167173-85-5, UK 2A

RL: BAC (Biological activity or effector, except adverse); PRP (Properties); BIOL (Biological study)

(UK-2A, B, C and D, novel antifungal antibiotics from Streptomyces sp. 517-02. I. Fermn., isolation, and biol. properties)

RN 167173-85-5

ANSWER 8 OF 9 HCAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER:

1995:934118 HCAPLUS

DOCUMENT NUMBER:

123:337552

TITLE:

Fungicides manufacture with Streptoverticillium

INVENTOR(S):

Taniguchi, Makoto; Shibata, Kozo; Abe, Keiichi;

Kodama, Tooru; Uotani, Kazumichi; Oonishi, Yoshitaka

PATENT ASSIGNEE(S):

Suntory Ltd, Japan; Meiji Seika Co

SOURCE:

Jpn. Kokai Tokkyo Koho, 10 pp.

CODEN: JKXXAF

DOCUMENT TYPE:

Patent

LANGUAGE:

Japanese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE JP 07233165 A2 19950905 JP 1994-26884 19940224 <--

OTHER SOURCE(S):

MARPAT 123:337552

GΙ

Fungicides (I: R = linear or branched aliph. (un)satd. acyl group) are AΒ manufd. by culturing Streptoverticillium sp. SAM2084. Shake-culture of Streptoverticillium sp. SAM2084 for manuf. of four I wherein R =2-methylpropanoyl (UK-2A), trans-2-methyl-2-butenoyl (UK-2B), 3-methylbutanoyl (UK-2C), and 2-methylbutanoyl (UK-2D) was shown. Also given were the physiol. and morphol. characteristics of the Streptoverticillium sp. SAM2084.

Ι

ΙT 167173-85-5P, UK 2A

> RL: BPN (Biosynthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (fungicides manuf. with Streptoverticillium)

RN 167173-85-5

ANSWER 9 OF 9 HCAPLUS COPYRIGHT 2001 ACS ACCESSION NUMBER: 1995:671786 HCAPLUS

DOCUMENT NUMBER:

123:164736

TITLE:

The structures of UK-1 and UK-2, novel antibiotics

from Streptomyces sp. 517-02

AUTHOR(S):

Hanafi, O Muhammad; Kozo, Shibata; Masaru, Kashiwada;

Masashi, Ueki; Makoto, Taniguchi

CORPORATE SOURCE:

Faculty Science, Osaka City University, Japan

SOURCE:

Tennen Yuki Kagobutsu Toronkai Koen Yoshishu (

1994), 36th, 728-35

CODEN: TYKYDS

DOCUMENT TYPE:

Journal Japanese

LANGUAGE: Jap

AB The mycelial cake was extd. with acetone, and filtered. The filtrate was concd. to give aq. soln., which was extd. with chloroform. Org. layer was

concd. to yield an oily material, followed by purifn. on silica gel column

chromatog. to give crude UK-1 and UK-2. Finally, the recrystn. of each fractions from MeOH, afforded UK-1 and UK-2. UK-1 (I), a novel metabolite, demonstrated potent cytotoxic activity against B16, Hela and P388 cells, and UK-2, novel complex of antibiotics, exhibited strong antifungal activity. Methylation of UK-1 by CH3I and anhyd. K2CO3 in dry acetone gave monomethyl ether deriv., Me-UK-1. Alk. hydrolysis of UK-1 afforded carboxylic acid deriv., DeMe-UK-1. Partial structures, A, B,

and

C were elucidated by COSY, and COLOC expts. Based on these results, the structure of UK-1 was deduced to be a novel benzoxazole dimer deriv. UK-2, novel metabolite contg. complex of antibiotics with strong antifungal activity, was purified by reverse phase HPLC, to give UK-2A,

В,

C and D. From NMR and mass spectral data, the structures of UK-2A, B, C and D were established as isobutyrate, tiglate, isovalerate, and 2-methylbutyrate of nine membered dilactone skeleton, resp. Based on the result of synthesis of hydrolysis products, the abs. configuration of

UK∸2

was identified.

IT 167173-85-5, Antibiotic UK 2A

RL: PRP (Properties)

(structures of UK-1 and UK-2, novel antibiotics from Streptomyces sp. 517-02)

RN 167173-85-5

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